4164-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket Nos. FDA-2020-D-1136, FDA-2020-D-1137, FDA-2020-D-1138, FDA-2020-D-1139]

Guidance Documents Related to Coronavirus Disease 2019 (COVID-19); Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of availability.

SUMMARY: The Food and Drug Administration (FDA or Agency) is announcing the availability of FDA guidance documents related to the Coronavirus Disease 2019 (COVID-19) public health emergency (PHE). This notice of availability (NOA) is pursuant to the process that FDA announced, in the *Federal Register* of March 25, 2020, for making available to the public COVID-19-related guidances. The guidances identified in this notice address issues related to the COVID-19 PHE and have been issued in accordance with the process announced in the March 25, 2020, notice. The guidances have been implemented without prior comment, but they remain subject to comment in accordance with the Agency's good guidance practices.

DATES: The announcement of the guidances is published in the *Federal Register* on [INSERT DATE OF PUBLICATION IN THE *FEDERAL REGISTER*]. The guidances have been implemented without prior comment, but they remain subject to comment in accordance with the Agency's good guidance practices.

ADDRESSES: You may submit either electronic or written comments on Agency guidances at any time as follows:

Electronic Submissions

Submit electronic comments in the following way:

- Federal eRulemaking Portal: https://www.regulations.gov. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to https://www.regulations.gov will be posted to the docket unchanged. Because your comment will be made public, you are solely responsible for ensuring that your comment does not include any confidential information that you or a third party may not wish to be posted, such as medical information, your or anyone else's Social Security number, or confidential business information, such as a manufacturing process. Please note that if you include your name, contact information, or other information that identifies you in the body of your comments, that information will be posted on https://www.regulations.gov.
- If you want to submit a comment with confidential information that you do not wish to be made available to the public, submit the comment as a written/paper submission and in the manner detailed (see "Written/Paper Submissions" and "Instructions").

Written/Paper Submissions

Submit written/paper submissions as follows:

- Mail/Hand delivery/Courier (for written/paper submissions): Dockets Management Staff (HFA-305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.
- For written/paper comments submitted to the Dockets Management Staff, FDA will post
 your comment, as well as any attachments, except for information submitted, marked and
 identified, as confidential, if submitted as detailed in "Instructions."

Instructions: All submissions received must include the name of the guidance(s) that the comments address and the docket number for the guidance (see table 1). Received comments

will be placed in the docket(s) and, except for those submitted as "Confidential Submissions," publicly viewable at https://www.regulations.gov or at the Dockets Management Staff between 9 a.m. and 4 p.m., Monday through Friday.

• Confidential Submissions--To submit a comment with confidential information that you do not wish to be made publicly available, submit your comments only as a written/paper submission. You should submit two copies total. One copy will include the information you claim to be confidential with a heading or cover note that states "THIS DOCUMENT CONTAINS CONFIDENTIAL INFORMATION." The Agency will review this copy, including the claimed confidential information, in its consideration of comments. The second copy, which will have the claimed confidential information redacted/blacked out, will be available for public viewing and posted on https://www.regulations.gov. Submit both copies to the Dockets Management Staff. If you do not wish your name and contact information to be made publicly available, you can provide this information on the cover sheet and not in the body of your comments and you must identify this information as "confidential." Any information marked as "confidential" will not be disclosed except in accordance with 21 CFR 10.20 and other applicable disclosure law. For more information about FDA's posting of comments to public dockets, see 80 FR 56469, September 18, 2015, or access the information at:

https://www.govinfo.gov/content/pkg/FR-2015-09-18/pdf/2015-23389.pdf.

Docket: For access to the docket to read background documents or the electronic and written/paper comments received, go to https://www.regulations.gov and insert the docket number, found in brackets in the heading of this document, into the "Search" box and follow the

prompts and/or go to the Dockets Management Staff, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

You may submit comments on any guidance at any time (see 21 CFR 10.115(g)(5)). Submit written requests for single copies of any of these guidances to the addresses noted in table 1. Send two self-addressed adhesive labels to assist that office in processing your requests. See the SUPPLEMENTARY INFORMATION section for electronic access to the guidance document.

FOR FURTHER INFORMATION CONTACT: Stephen Ripley, Center for Biologics

Evaluation and Research (CBER), Food and Drug Administration, 10903 New Hampshire Ave.,

Bldg. 71, Rm. 7268, Silver Spring, MD 20993-0002, 240-402-7911; Erica Takai, Center for

Devices and Radiological Health (CDRH), Food and Drug Administration, 10903 New

Hampshire Ave., Bldg. 66, Rm. 5456, Silver Spring, MD 20993-0002, 301-796-6353; Kimberly

Thomas, Center for Drug Evaluation and Research (CDER), Food and Drug Administration,

10903 New Hampshire Ave., Bldg. 51, Rm. 6220, Silver Spring, MD 20993-0002, 301-796
2357; Phil Chao, Center for Food Safety and Applied Nutrition (CFSAN), CPK1 Rm 1C001,

HFS-024, Food and Drug Administration, College Park, MD 20740, 240-402-2112.

SUPPLEMENTARY INFORMATION:

I. Background

On January 31, 2020, as a result of confirmed cases of COVID-19, and after consultation with public health officials as necessary, Alex M. Azar II, Secretary of Health and Human Services, pursuant to the authority under section 319 of the Public Health Service Act (42 U.S.C. 247d) (PHS Act), determined that a PHE exists and has existed since January 27, 2020,

nationwide.¹ On March 13, 2020, President Donald J. Trump declared that the COVID-19 outbreak in the United States constitutes a national emergency, beginning March 1, 2020.²

In the Federal Register of March 25, 2020 (the March 25, 2020, notice) (available at: https://www.govinfo.gov/content/pkg/FR-2020-03-25/pdf/2020-06222.pdf), FDA announced procedures for making available FDA guidances related to the COVID-19 PHE. These procedures, which operate within FDA's established good guidance practices regulations, are intended to allow FDA to rapidly disseminate Agency recommendations and policies related to COVID-19 to industry, FDA staff, and other stakeholders. The March 25, 2020, notice stated that due to the need to act quickly and efficiently to respond to the COVID-19 PHE, FDA believes that prior public participation will not be feasible or appropriate before FDA implements COVID-19-related guidances. Therefore, FDA will issue COVID-19-related guidances for immediate implementation without prior public comment (see section 701(h)(1)(C) of the Federal Food, Drug, and Cosmetic Act (FD&C Act) (21 U.S.C. 371(h)(1)(C) and 21 CFR 10.115(g)(2) (§ 10.115(g)(2))). The guidances are available at FDA's webpage entitled "COVID-19-Related Guidance Documents for Industry, FDA Staff, and Other Stakeholders" (https://www.fda.gov/emergency-preparedness-and-response/mcm-issues/covid-19-relatedguidance-documents-industry-fda-staff-and-other-stakeholders) and through FDA's webpage entitled "Search for FDA Guidance Documents" available at https://www.fda.gov/regulatoryinformation/search-fda-guidance-documents.

The March 25, 2020, notice further stated that, in general, rather than publishing a separate NOA for each COVID-19-related guidance, FDA intends to publish periodically a

¹ On April 21, 2020, the PHE Determination was extended, effective April 26, 2020. These PHE Determinations are available at https://www.phe.gov/emergency/news/healthactions/phe/Pages/default.aspx.

² Proclamation on Declaring a National Emergency Concerning the Novel Coronavirus Disease (COVID-19) Outbreak (March 13, 2020), available at https://www.whitehouse.gov/presidential-actions/proclamation-declaring-national-emergency-concerning-novel-coronavirus-disease-covid-19-outbreak/.

consolidated NOA announcing the availability of certain COVID-19-related guidances FDA issued during the relevant period, as included in table 1. This notice announces COVID-19-related guidances that are posted on FDA's website.

II. Availability of COVID-19-Related Guidances

Pursuant to the process described in the March 25, 2020, notice, FDA is announcing the availability of the following COVID-19-related guidances:

Table 1.--Guidances Related to the COVID-19 Public Health Emergency

Docket No.	Center/Office	Title of Guidance	Contact Information to Request
			Single Copies
FDA-2020-D-	CBER	Investigatory COVID-19 Convalescent	Office of Communication, Outreach
1137		Plasma (April 2020) (Updated May 1, 2020)	and Development, 10903 New
			Hampshire Ave., Bldg. 71, Rm.
			3128, Silver Spring, MD 20993-
			0002, 1-800-835-4709 or 240-402-
			8010, email ocod@fda.hhs.gov
FDA-2020-D-	CDRH	Enforcement Policy for Clinical Electronic	CDRH-Guidance@fda.hhs.gov
1138		Thermometers During the Coronavirus	Please include the document number
		Disease 2019 (COVID-19) Public Health	20014 and complete title of the
		Emergency (April 4, 2020)	guidance in the request.
FDA-2020-D-	CDRH	Enforcement Policy for Infusion Pumps and	CDRH-Guidance@fda.hhs.gov
1138		Accessories During the Coronavirus Disease	Please include the document number
		2019 (COVID-19) Public Health Emergency	20014 and complete title of the
		(April 5, 2020)	guidance in the request.
FDA-2020-D-	CDRH	Enforcement Policy for Remote Ophthalmic	CDRH-Guidance@fda.hhs.gov
1138		Assessment and Monitoring Devices During	Please include the document number
		the Coronavirus Disease 2019 (COVID-19)	20014 and complete title of the
		Public Health Emergency (April 6, 2020)	guidance in the request.
FDA-2020-D-	CDRH	Enforcement Policy for Extracorporeal	CDRH-Guidance@fda.hhs.gov
1138		Membrane Oxygenation and	Please include the document number
		Cardiopulmonary Bypass Devices During	20014 and complete title of the
		the Coronavirus Disease 2019 (COVID-19)	guidance in the request.
		Public Health Emergency (April 6, 2020)	
FDA-2020-D-	CDRH	Enforcement Policy for Digital Health	CDRH-Guidance@fda.hhs.gov
1138		Devices for Treating Psychological	Please include the document number
		Disorders During the Coronavirus Disease	20014 and complete title of the
		2019 (COVID-19) Public Health Emergency	guidance in the request.
		(April 14, 2020)	
FDA-2020-D-	CDRH	Enforcement Policy for Telethermographic	CDRH-Guidance@fda.hhs.gov
1138		Systems During the Coronavirus Disease	Please include the document number
		2019 (COVID-19) Public Health Emergency	20014 and complete title of the
		(April 16, 2020)	guidance in the request.
FDA-2020-D-	CDRH	Enforcement Policy for Non-Invasive Fetal	CDRH-Guidance@fda.hhs.gov
1138		and Maternal Monitoring Devices Used to	Please include the document number
		Support Patient Monitoring During the	20014 and complete title of the
		Coronavirus Disease 2019 (COVID-19)	guidance in the request.
		Public Health Emergency (April 23, 2020)	
FDA-2020-D-	CDRH	Enforcement Policy for Imaging Systems	CDRH-Guidance@fda.hhs.gov

1138		During the Coronavirus Disease 2019 (COVID-19) Public Health Emergency (April 23, 2020)	Please include the document number 20014 and complete title of the guidance in the request.
FDA-2020-D- 1138	CDRH	Enforcement Policy for Remote Digital Pathology Devices During the Coronavirus Disease 2019 (COVID-19) Public Health Emergency (April 24, 2020)	CDRH-Guidance@fda.hhs.gov Please include the document number 20014 and complete title of the guidance in the request.
FDA-2020-D- 1136	CDER	Temporary Policy Regarding Non-Standard PPE Practices for Sterile Compounding by Pharmacy Compounders not Registered as Outsourcing Facilities During the COVID-19 Public Health Emergency (April 10, 2020)	druginfo@fda.hhs.gov Please include the docket number FDA-2020-D-1136 and complete title of the guidance in the request.
FDA-2020-D- 1136	CDER	Policy for the Temporary Use of Portable Cryogenic Containers Not in Compliance With 21 CFR 211.94(e)(1) For Oxygen and Nitrogen During the COVID-19 Public Health Emergency (April 2020) (Updated April 20, 2020)	druginfo@fda.hhs.gov Please include the docket number FDA-2020-D-1136 and complete title of the guidance in the request.
FDA-2020-D- 1136	CDER	Temporary Policy for Compounding of Certain Drugs for Hospitalized Patients by Outsourcing Facilities During the COVID- 19 Public Health Emergency (April 16, 2020) (Updated May 8, 2020)	druginfo@fda.hhs.gov Please include the docket number FDA-2020-D-1136 and complete title of the guidance in the request.
FDA-2020-D- 1136	CDER	Temporary Policy for Compounding of Certain Drugs for Hospitalized Patients by Pharmacy Compounders not Registered as Outsourcing Facilities During the COVID- 19 Public Health Emergency Guidance for Industry (April 20, 2020) (Updated May 8, 2020)	druginfo@fda.hhs.gov Please include the docket number FDA-2020-D-1136 and complete title of the guidance in the request.
FDA-2020-D- 1136	CDER	Temporary Policy on Repackaging or Combining Propofol Drug Products During the COVID-19 Public Health Emergency (April 22, 2020)	druginfo@fda.hhs.gov Please include the docket number FDA-2020-D-1136 and complete title of the guidance in the request.
FDA-2020-D- 1139	CFSAN	Temporary Policy on Regulatory Enforcement of 21 CFR Part 118 (the Egg Safety Rule) During the COVID-19 Public Health Emergency (April 6, 2020)	Office of Nutrition and Food Labeling, Food Labeling and Standards Staff, Center for Food Safety and Applied Nutrition, Food and Drug Administration, 5001 Campus Dr., College Park, MD 20740

Although these guidances have been implemented immediately without prior comment, FDA will consider all comments received and revise the guidances as appropriate (see § 10.115(g)(3)).

These guidances are being issued consistent with FDA's good guidance practices regulation (§ 10.115). The guidances represent the current thinking of FDA. They do not

establish any rights for any person and are not binding on FDA or the public. You can use an alternative approach if it satisfies the requirements of the applicable statutes and regulations.

III. Paperwork Reduction Act of 1995

A. CBER

The guidance indicated below refers to previously approved collections of information. These collections of information are subject to review by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995 (44 U.S.C. 3501-3521) (PRA). The collections of information in the following FDA regulations and guidance have been approved by OMB as listed in the following table:

Table 2.--CBER Guidance

COVID-19 Guidance	CFR Cite Referenced in	Another Guidance Title	OMB Control
Title	COVID-19 Guidance	Referenced in COVID-19	No(s).
		Guidance	
Investigatory COVID-19	21 CFR part 312	N/A	0910-0014
Convalescent Plasma	21 CFR 606.121		0910-0116
	21 CFR part 630		0910-0116
	Form FDA 3926		0910-0814

B. CDRH

The guidances listed below refer to previously approved collections of information.

These collections of information are subject to review by OMB under the PRA. The collections of information in the following FDA regulations and guidance have been approved by OMB as listed in the following table:

Table 3.--CDRH Guidances

COVID-19 Guidance Title	CFR Cite Referenced in	Another Guidance Title	OMB Control
COVID 1) Guidance Tric	COVID-19 Guidance	Referenced in COVID-	No(s).
	COVID 17 Guidance	19 Guidance	110(5).
Enforcement Policy for Clinical	21 CFR part 807, subpart E		0910-0120
Electronic Thermometers During	21 CFR part 806		0910-0359
the Coronavirus Disease 2019	21 CFR part 807, subparts A		
(COVID-19) Public Health	through D		0910-0625
Emergency	21 CFR parts 830 & 801.20		0910-0720
	21 CFR parts 800, 801, 809		0910-0485
	21 CFR part 820		0910-0073
Enforcement Policy for Infusion		Emergency Use	
Pumps and Accessories During the		Authorization of	
Coronavirus Disease 2019		Medical Products and	
(COVID-19) Public Health		Related Authorities;	
Emergency		Guidance for Industry	
		and Other Stakeholders	0910-0595
	21 CFR part 807, subpart E		0910-0120
	21 CFR part 807, subparts A		
	through D		0910-0625
	21 CFR part 820		0910-0073
	21 CFR part 806		0910-0359
	21 CFR parts 830 and 801.20		0910-0720
	21 CFR parts 800, 801, and 809		0910-0485
	21 CFR part 803		0910-0437
Enforcement Policy for Remote	21 CFR part 807, subpart E		0910-0120
Ophthalmic Assessment and	21 CFR part 807, subparts A		
Monitoring Devices During the	through D		0910-0625
Coronavirus Disease 2019	21 CFR part 822		0910-0449
(COVID-19) Public Health	21 CFR part 820		0910-0073
Emergency	21 CFR part 806		0910-0359
	21 CFR parts 830 and 801.20		0910-0720
Enforcement Daling for	21 CFR parts 800, 801, and 809	E	0910-0485
Enforcement Policy for		Emergency Use Authorization of	
Extracorporeal Membrane Oxygenation and		Medical Products and	
Cardiopulmonary Bypass Devices		Related Authorities;	
During the Coronavirus Disease		Guidance for Industry	
2019 (COVID-19) Public Health		and Other Stakeholders	0910-0595
Emergency	21 CFR part 807, subpart E	and Other Stakeholders	0910-0393
Lineigency	21 CFR part 814, subparts A		0710-0120
	through E		0910-0231
	21 CFR parts 800, 801, and 809		0910-0485
	21 CFR part 820		0910-0073
	21 CFR part 803		0910-0437
Enforcement Policy for Digital	21 CFR part 807, subpart E		0910-0120
Health Devices for Treating	21 CFR part 806		0910-0359
Psychiatric Disorders During the	21 CFR part 807, subparts A		
Coronavirus Disease 2019	through D		0910-0625
(COVID-19) Public Health	21 CFR parts 830 and 801.20		0910-0720
Emergency	21 CFR parts 800, 801, and 809		0910-0485

		1	
Enforcement Policy for	21 CFR parts 800, 801, and 809		0910-0485
Telethermographic Systems	21 CFR part 806		0910-0359
During the Coronavirus Disease	21 CFR part 807, subparts A		
2019 (COVID-19) Public Health	through D		0910-0625
Emergency	21 CFR part 807, subpart E		0910-0120
	21 CFR part 820		0910-0073
	21 CFR part 822		0910-0449
	21 CFR parts 830 and 801.20		0910-0720
Enforcement Policy for Non-	21 CFR part 807, subpart E		0910-0120
Invasive Fetal and Maternal	21 CFR parts 800, 801, and 809		0910-0485
Monitoring Devices Used to	21 CFR part 820		0910-0073
Support Patient Monitoring During			
the Coronavirus Disease 2019			
(COVID-19) Public Health			
Emergency			
Enforcement Policy for Imaging	21 CFR parts 800, 801, and 809		0910-0485
Systems During the Coronavirus	21 CFR part 807, subpart E		0910-0120
Disease 2019 (COVID-19) Public	21 CFR part 814, subparts A		0,10 0120
Health Emergency	through E		0910-0231
Treatin Emergency	21 CFR part 820		0910-0073
	21 CFR parts 1000-1050		0910-0025
Enforcement Policy for Remote	21 61 11 parts 1000 1000	Administrative	0310 0023
Digital Pathology Devices During		Procedures for CLIA	
the Coronavirus Disease 2019		Categorization:	
(COVID-19) Public Health		Guidance for Industry	
Emergency		and Food and Drug	
Zinergene j		Administration Staff	0910-0667
	21 CFR part 807, subpart E	Tammoutuon Sum	0910-0120
	21 CFR part 812		0910-0078
	21 CFR part 820		0910-0073
	21 CFR part 820 21 CFR parts 830 and 801.20		0910-0720
	21 CFR parts 800, 801, and 809		0910-0485
	21 CFR part 814, subpart H		0910-0332
	21 CFR part 814, subpart 11 21 CFR part 820		0910-0332
	21 CFR parts 800, 801, and 809		0910-0073
	21 C1 K parts 600, 601, and 609		0710-0463

C. CDER

The guidances listed below refer to previously approved collections of information.

These collections of information are subject to review by OMB under the PRA. The collections of information in the following FDA regulations and guidances have been approved by OMB as listed in the following table:

Table 4.--Guidances and Regulations

GOVERN 10 G 11 FE11	Table 4Guidances and		0.00
COVID-19 Guidance Title	CFR or FD&C Act Cite Referenced in COVID-19 Guidance	Another Guidance Title Referenced in COVID-19 Guidance	OMB Control No(s).
Policy for Temporary Use of Portable Cryogenic Containers Not in Compliance With 21 CFR 211.94 for Oxygen and Nitrogen During COVID-19 Public Health Emergency	21 CFR parts 201, 210, 211.84, 211.94, and 211.100	Current Good Manufacturing Practice for Medical Gases Medical Gas Containers and Closures; Current Good Manufacturing Practice Requirements	0910-0139
Temporary Policy for	21 CFR 314.81	Current Good Manufacturing	0910-0777
Compounding of Certain Drugs	21 CFR 600.82	PracticeGuidance for Human Drug	0910-0338
for Hospitalized Patients by	Section $503B(b)(1)(A)(i)$ of	Compounding Outsourcing	0910-0001
Outsourcing Facilities During the COVID-19 Public Health Emergency	the FD&C Act (21 U.S.C. 353b(b)(1)(A)(i)	Facilities Under Section 503B of the FD&C Act	0910-0139
Temporary Policy for		Compounded Drug Products That	0910-0001
Compounding of Certain Drugs		are Essentially Copies of a	0910-0139
for Hospitalized Patients by		Commercially Available Drug	0910-0338
Pharmacy Compounders not Registered as Outsourcing		Product under Section 503A of the Federal Food, Drug and Cosmetic	
Facilities During the COVID-19		Act	
Public Health Emergency		Temporary Policy for Compounding	
		of Certain Drugs for Hospitalized	
		Patients by Outsourcing Facilities	
		During the COVID-19 Public	
		Health Emergency	
		Prescription Requirement Under	
		Section 503A of the Federal Food, Drug, and Cosmetic Act	
		Temporary Policy Regarding Non-	
		Standard PPE Practices for Sterile	
		Compounding by Pharmacy	
		Compounders not Registered as	
		Outsourcing Facilities during the	
		COVID-19 Public Health	
		Emergency	0010 0120
Temporary Policy on		Repackaging of Certain Human	0910-0139
Repackaging or Combining Propofol Drug Products During		Drugs by Pharmacies and Outsourcing Facilities	0910-0572 0910-0777
the COVID-19 Public Health		Temporary Policy for Compounding	0910-0777
Emergency		of Certain Drugs for Hospitalized	3710 0000
,-8,		Patients by Pharmacy Compounders	
		not Registered as Outsourcing	
		Facilities During the COVID-19	
		Public Health Emergency	
		Temporary Policy for Compounding	
		of Certain Drugs for Hospitalized Patients by Outsourcing Facilities	
		During the COVID-19 Public	
		Health Emergency	
	l	Treatm Differency	

The guidance indicated below refers to previously approved collections of information.

These collections of information are subject to review by OMB under the PRA. The collections

of information in the following FDA regulations and guidance have been approved by OMB as listed in the below table. This guidance also contains a new collection of information not approved under a current collection. This new collection of information has been granted a PHE waiver from the PRA by HHS on March 19, 2020, under section 319(f) of the PHS Act. Information concerning the PHE PRA waiver can be found on the HHS website at https://aspe.hhs.gov/public-health-emergency-declaration-pra-waivers.

Table 5.--New PRA Information Collection

COVID-19 Guidance Title	CFR Cite Referenced in COVID-19 Guidance	Another Guidance Referenced in COVID-19 Guidance	OMB Control No.	New Collection Covered by PHE PRA Waiver
Temporary Policy Regarding Non- Standard PPE Practices for Sterile Compounding by Pharmacy Compounders not Registered as Outsourcing Facilities During the COVID-19 Public Health Emergency	21 CFR parts 210 and 211	Enforcement Policy for Face Masks and Respirators During the Coronavirus Disease (COVID-19) Public Health Emergency (Revised) Enforcement Policy for Gowns, Other Apparel, and Gloves During the Coronavirus Disease (COVID-19) Public Health Emergency Electronic Drug Product Reporting for Human Drug Compounding Outsourcing Facilities Under Section 503B of the Federal Food, Drug, and Cosmetic Act	0910-0139	Recordkeeping of compounding without standard PPE; recordkeeping of any change of sterilization/aseptic processing methods; documentation of mitigation strategies for sterile compounding without standard PPE

D. CFSAN

The guidance indicated below refers to previously approved collections of information.

These collections of information are subject to review by OMB under the PRA. The collections of information in the following FDA regulations and guidance have been approved by OMB as listed in the following table:

Table 6.--CFSAN Guidance

COVID-19 Guidance Title	CFR Cite Referenced	Another Guidance Title	OMB
	in COVID-19	Referenced in COVID-	Control No.
	Guidance	19 Guidance	
Temporary Policy	21 CFR part 118		0910-0660
Regarding Enforcement of			
21 CFR Part 118 (the Egg			
Safety Rule) During the			
COVID-19 Public Health			
Emergency			

IV. Electronic Access

Persons with access to the internet may obtain COVID-19-related guidances at:

- the FDA webpage entitled "COVID-19-Related Guidance Documents for Industry, FDA Staff, and Other Stakeholders," available at https://www.fda.gov/emergency-preparedness-and-response/mcm-issues/covid-19-related-guidance-documents-industry-fda-staff-and-other-stakeholders;
- the FDA webpage entitled "Search for FDA Guidance Documents," available at https://www.fda.gov/regulatory-information/search-fda-guidance-documents; or
- https://www.regulations.gov.

Dated: May 19, 2020.

Lowell J. Schiller,

Principal Associate Commissioner for Policy.

[FR Doc. 2020-11238 Filed: 5/22/2020 8:45 am; Publication Date: 5/26/2020]